		Tes	t Device	Tempera	ature Lo	g		
Testing S	ite:				City:		_	
Testing K	its Location:							
Type of R	Rapid Test Kits:	□ Determin	ne 🗆 Insti 🏻	☐ Syphilis He	alth Check	□ OraQuicl	k □ Uni-Ge	old Clearvie
temperatu between o	and low tempera are range memor checks. emperature fall	y that will dis	play the warm	est and cooles	t temperature	s reached in th	ne storage area	in-
1	Allowable Temp	Range:	from: de	egrees F	to:	degrees F		
Daily Temperature Record for Month:Year:								
Date	Low	High	Initial	Date	Low	High	Initial	
1				16				
2				17				
3				18				
4				19				
5				20				
6				21				
7				22				
8				23				
9				24				
10				25				
11				26				
12				27				
13				28				
14				29				
15				30				
l				31				i
Note on	y incidents aı	nd correctiv	zo octions tol	zon holow:				
Note an	y meidents ai	iu correctiv		ective Actio	n			
Date:								
Quality A	Assurance Coord	inator				D	ate:	

Attachment RT-3.2 (maintain on-site) Control Kit Temperature Log									
		<u>Co</u>	ntroi Kit	ı empera	iture Log	J			
Testing Site:	Testing Site: City:								
Control Kits	location:								
Type of Rapid Test Control Kits: ☐ Determine ☐ Insti ☐ Syphilis Health Check ☐ OraQuick ☐ Uni-Gold ☐ Clearview									
The high and low temperatures of the control kit storage refrigerator should be recorded using a digital thermometer with a temperature range memory that will display the warmest and coolest temperatures reached in the refrigerator in between checks. If temperature falls outside the allowable range, notify quality assurance coordinator immediately.									
Alle	owable Temp	Range:	from: de	grees F	to: o	legrees F			
	Daily	Temperatur	e Record for N	Month:	Y	ear:	_		
Date	Low	High	Initial	Date	Low	High	Initial	1	
1				16					
2				17					
3				18					
4				19					
5				20					
6				21					
7				22					
8				23					
9				24					
10				25					
11				26					
12				27					
13				28					
14				29					
15				30					
				31					
Note any incidents and corrective actions taken below: Corrective Action									
date:			Corre	CHYC ACHU	11				
Quality Assurance Coordinator Date:									

Attachment RT-3.3 (maintain o	n-site)					
	Daily Rapid HIV Test Log					
Test Site:	Date of Testing:					
(note the lot number from the test kit package, not the outer box or shipment materials)						
Types of Rapid Test:	Determine, Insti, Syphilis Health Check, OraQuick, Uni-Gold, Clearview					

Type of Rapid Test	Rapid Lab Counselor #	HIV Test form Number	Room Temperature	Time Test Started	Time Test Result Read	Rapid Test Result	Date Client Notified	Lot Number of Test Kit	Test Kit Expiration Date
						☐ Reactive			
						\square Ag \square Ab			
						☐ Invalid			
						☐ Reactive			
						\square Ag \square Ab			
						☐ Invalid			
						☐ Reactive			
						\Box Ag \Box Ab			
						□ Neg			
						□ Invalid			
						☐ Reactive			
						□ Ag □ Ab			
						□ Neg			
						□ Invalid			
						□ Reactive			
						□ Ag □ Ab			
						□ Neg			
						□ Invalid			
						☐ Reactive			
						\Box Ag \Box Ab			
						□ Neg			
						☐ Invalid			

Quality Assurance Coordinator:	Date:	
- ·		

Control Kit Log									
Test Site:	Test Site: Month/Year:								
Control I				Ma	Manufacturer's Expiration Date:				
Date Kits	Date Kits Opened:								
Type of Kit	Controls: D	etermine, Insti, S	yphilis He	alth Check,	OraQuick,	Uni-Gold, C	learview		
Type of Kit			MEG	11117 1	1111/2		G 1.11	Reason for running	
Controls	Date	Counselor #	NEG	HIV-1	HIV-2	Antigen	Syphilis	controls	
			☐ Pass ☐ Fail	□ Pass □ Fail	☐ Pass ☐ Fail	□ Pass □ Fail	□ Pass □ Fail		
			☐ Pass	☐ Pass	□ Pass	□ Pass	□ Pass		
			☐ Fail	☐ Fail	☐ Fail	☐ Fail	☐ Fail		
			□ Pass	Pass	Pass	Pass	□ Pass		
			☐ Fail	☐ Fail	☐ Fail	☐ Fail	☐ Fail		
			☐ Pass ☐ Fail	□ Pass □ Fail	☐ Pass ☐ Fail	☐ Pass ☐ Fail	☐ Pass ☐ Fail		
				□ Pass	□ Pass	Pass	Pass		
			☐ Fass	□ Fail	□ Fass	☐ Fass	□ Fail		
			□ Pass	□ Pass	□ Pass	Pass	□ Pass		
			☐ Fail	☐ Fail	□ Fail	□ Fail	□ Fail		
			□ Pass	□ Pass	□ Pass	□ Pass	□ Pass		
			□ Fail	□ Fail	□ Fail	□ Fail	□ Fail		
			☐ Pass	☐ Pass	□ Pass	□ Pass	□ Pass		
			☐ Fail	☐ Fail	☐ Fail	☐ Fail	☐ Fail		
			☐ Pass	☐ Pass	☐ Pass	☐ Pass	☐ Pass		
			☐ Fail	☐ Fail	☐ Fail	☐ Fail	☐ Fail		
			☐ Pass	☐ Pass	☐ Pass	□ Pass	□ Pass		
			☐ Fail	☐ Fail	☐ Fail	☐ Fail	□ Fail		
			□ Pass	☐ Pass	□ Pass	□ Pass	□ Pass		
			☐ Fail	☐ Fail	☐ Fail	☐ Fail	☐ Fail		
			□ Pass	□ Pass	□ Pass	□ Pass	□ Pass		
			☐ Fail	☐ Fail	☐ Fail	☐ Fail	☐ Fail		
			□ Pass	□ Pass	Pass	Pass	□ Pass		
			☐ Fail	☐ Fail	☐ Fail	☐ Fail	☐ Fail		
			☐ Pass	☐ Pass	□ Pass	☐ Pass	☐ Pass		
			☐ Fail	☐ Fail	☐ Fail	☐ Fail	☐ Fail		
			☐ Pass ☐ Fail	☐ Pass ☐ Fail	☐ Pass ☐ Fail	☐ Pass ☐ Fail	☐ Pass ☐ Fail		
1			L L'all	□ 1 an	– 1 an	□ 1 an	– 1 (111		
Quality Ass	urance Coord	linator:					Date:		

Possible reasons for running controls: New Shipment, New Lot Number, Storage or Operating Temperature Out of Range, Arrived at Outreach Location, Facility Protocol

Attachment RT-3.5 (submit to SHP monthly)

HIV AND SYPE	HIV AND SYPHILIS TESTING SUPPLY ORDER FORM							
Contact Information (Agency condu	cting HIV Testin	ng):						
Testing Site Name:	Testing Site Name: Order Date:							
Quality Assurance Coordinator:								
Mailing Address:								
City, State, Zip:			hone Number:					
Fax Number:			ess:					
CLIA Certificate #:(Req								
Please write the number of cases/bordelivery or pick up. Some items may Agencies located within Region 1 will LIST OF SUPPLIES	not be available not be notified whe	at the time of order	or available to your site. ly for pick up.					
HIV Test forms-Part 1	100 forms/packe	et						
Sites must have prior approval from (OPH SHP before	ordering any of the f	following items:					
Determine Ag/Ab Rapid Test Kits Determine Ag/Ab Kit Control Determine Fingerstick Accessory Kit INSTI Rapid Test Kits INSTI Kit Control Syphilis Health Check Rapid Test Kits Syphilis Health Check Kit Control Digital Memory Thermometer Timer XL Gloves □ Nitrile □ Latex L Gloves □ Nitrile □ Latex M Gloves □ Nitrile □ Latex S Gloves □ Nitrile □ Latex Workspace Covers Biohazard waste disposal bag Sharps Container (limited availability) Please fax this form to: OPH SHP PU	25 kits/box 1 kit/box 1 kit/box 50 kits/box 1 kit/box 20 kits/box 1 kit/box Each Each 100/box 100/box 100/box 100/box 100/box Each Each Each	PPLIES COORDINA						
For SHP Use Only:								
SHP Staff Initials:								
Rapid Tests Lot #:			n date:					
Control Lot #:		Control kit expiration date:						
Delivered to (name):		Date delivered:						

Attachment RT-3.6 (to be completed by Regional Coordinator and submitted as needed)

HIV Prevention Counseling, Testing and Referral (CTR) Rapid Site Assessment and Registration Form

All sites, whether fixed or mobile, must be registered with OPH SHP. Please allow four (4) weeks for processing.

Type of Request (check one)): ☐ New Site ☐ Update Existing Site ☐ Drop Site
Contact Information (Agency co	onducting CTR):
	Parish:
Phone Number:	Fax Number:
E-Mail Address:	CLIA Certificate #:
Is this agency conducting HIV test	ts as a part of the new CDC initiative? Yes No
Executive Director Information:	
Name:	
Mailing Address:	
City, State, Zip:	
Phone Number:	Fax Number:
Executive Director's Email:	
Prevention Manager Informatio	n•
G	
	Fax Number:
Quality Assurance Coordinator	Information:
Name:	
Mailing Address:	
City, State, Zip:	
Phone Number:	Fax Number:
Quality Assurance Coordinator's I	Email:

Site Information (location	where CTR will be conduc	cted):						
Name of Site:								
Site Address:								
City, State, Zip:								
Phone Number: Fax Number:								
Detailed Description of Site	Type (i.e. clientele, hours of	of operation, services offered):						
_	_	dentiality be assured, where in						
Type of Testing Requested ((check all that apply):							
☐ Rapid Testing:	OraSure	□ Blood (lab)						
Date:	Observed by:							
Check appropriate assessn	nent of testing site:							
Work space to process test:	☐ Acceptable ☐ Conditio	nal (describe) □ Unacceptable						
Confidential setting:	☐ Acceptable ☐ Conditio	nal (describe) □ Unacceptable						
Cleanliness:	☐ Acceptable ☐ Conditio	nal (describe) □ Unacceptable						
Lighting:	☐ Acceptable ☐ Conditio	nal (describe) □ Unacceptable						
Temperature control:	*	nal (describe) □ Unacceptable						
Supply storage:	-	nal (describe) □ Unacceptable						
Hand washing station:		nal (describe) □ Unacceptable						
Record keeping: Waiting area:	-	nal (describe) □ Unacceptable nal (describe) □ Unacceptable						
Č	•	,						
For Office Use Only: Date re-	quest received:	Date visited:						
Recommendation:								
SHP Coordinator Initials:	CTR Supervisor's Initials:	Date logged into database:						
Approved for: □ Rapid Testing:	Primary Test	Second Test	☐ Syphilis					
Health Check ☐ Syphilis Blood (lab) Site #:	Parent Site #:						

Attachment RT-3.8 (submit to SHP as needed)

Quality Assurance Coordinator Registration/Designation Form

All Agencies conducting Rapid HIV Testing in Louisiana must designate and register a Quality Assurance Coordinator. The Quality Assurance Coordinator should be a person with significant experience conducting rapid testing (6 months experience and a minimum of 200 rapid tests conducted) and familiar with storage and operating procedures/requirements of the rapid testing device(s) used at their agency.

<u>Submit to HAP immediately whenever the designated Quality Assurance Coordinator changes or when updates/changes to his/her contact information occur.</u>

Rapid Testing Site:	Site Number:						
Date Form Submitted:	Submitter:						
	_Newly Designated Quality Assurance Coordinator _Change in Quality Assurance Coordinator's contact information _Other, specify below:						
About the Designated	Quality Assurance Coordinator:						
Name*: Title*: Work Address*:							
Counselor Number*: Work Phone*: Cell: Alternate Phone Work Email*: Alternate Email:							
Number of Months/Ye	Number of Months/Years Experience with Rapid Testing:						
*#	reas marked with an asterisk are required fields						

Fax completed form to (504) 568-7044 Attention CTR Supervisor

Steps to HIV Counseling and Testing Certification

Steps for Obtaining a Counselor Number:

- 1. Attend a combined HIV & Syphilis Prevention Counseling and Rapid Testing course in its entirety and leave with a certificate of participation.
- 2. After completing the HIV & Syphilis Prevention Counseling and Rapid Testing training and receiving a certificate of completion, there are two additional steps. First, a written test covering HIV & syphilis prevention counseling, rapid testing skills, and protocol/paperwork must be passed. The dates, locations, and method of signing up for a class are outlined on www.hiv411.org. Secondly, all persons conducting CRT must successfully complete an observation session with the Regional Prevention Coordinator or other SHP Prevention staff as arranged by the Prevention Coordinator. Each person has two opportunities to pass the written test and the counselor observation. If the person fails either the test or the observation twice, they must go through the entire process again, beginning with training. Also, the written test must be passed before the observation can be scheduled.
- 3. Once the SHP Training Coordinator assigns a unique counselor number to the counselor, they are fully certified and may conduct CRT.

Steps for Registering a Rapid Testing Site:

- 1. Regional HIV Coordinator must conduct a site visit and make their recommendation on the site assessment and registration form. This form will then be given to the CT Supervisor.
- 2. If the site is favorably observed, CT Supervisor will assign a site number and mail a certificate with this number on it. A copy of this certificate must be kept on the site premises at all times.

Please Note: Meeting all counselor requirements does not automatically qualify your agency for site approval. Meeting all site requirements does not automatically qualify your agency for funding or free testing materials.

Attachment RT-3.10 (maintain on site-for information only)

Louisiana HIV Prevention Counseling and Rapid Testing Service Delivery Model

Step 1a - Introduce and Orient the Client to the Session

- Introduce yourself to the client.
- Assess client's readiness to receive the results on the same day.
- Offer options for testing (conventional or rapid) including HIV & syphilis.
- Describe the testing process, what type of specimen will be collected, how long the whole process will take, and what each of the three possible results mean.
- Explain to client that if a preliminary positive result is received, a confirmatory test should be conducted. The only exception is if Determine shows an antigen only positive.
- Address Partner Services, including informing the client that if results come back positive, a DIS will contact
 them to offer additional services.
- Offer anonymous and confidential options, and explain what each mean.
- Obtain Informed Consent.
- Provide appropriate subject information pamphlet for the rapid test being conducted.

Step 1b - Administer the Rapid Test

- Follow applicable universal precautions
- Clearly label the test device being used
- Demonstrate/facilitate specimen collection
- Start Timer

Step 2 – Identify Risk Behaviors and Circumstances

- Engage client in a discussion of risk behavior
- Assess client's previous experience with HIV testing and knowledge about HIV/AIDS
- Complete all but results section of HIV Test Form-Part 1

Step 3a – Identify Safer Goal Behaviors

- Give client information on relevant risk and harm reduction strategies
- Use relevant information pamphlets, brochures and/or brief videos
- Have client explain what he/she can do to reduce risk
- Assessing client readiness to receive results can continue up until the timer goes off
- Allow time for client to process and respond

Step 3b – Interpret and Deliver the Test Result (after appropriate time as elapsed)

- Follow applicable universal precautions for handling rapid testing materials
- Interpret Test Result (use a second reviewer if needed and client is not present)
- Return to client and give the results immediately in a simple and direct fashion
- Allow time for client to process and respond

Step 4 – Develop Risk Reduction/Action Plan (can be initiated prior to delivery of test results but should be modified, as needed, after results are provided)

- Based on the results of the test and the client's risk profile, assist the client in developing an action plan to further protect their health and the health of their partners.
- Document risk reduction plan in client's file

Step 5 – Offer Referrals and Provide Support (can be initiated prior to delivery of test results but should be modified, as needed, after results are provided)

Make appropriate referrals and negotiate plans to follow up with the client

Step 6 – Summarize and Close the Session

	1 (maintain on site-for info					1 m - 1		
	a Office of Public He							
	vention counselors an n prior to performing							
	fter and copies of all o							рег
Name of	_	Date		Counselor			Point Sca	ale:
Counselor:		Trained:	1	#:			0 = not do	
Date and Time of Observation:			Location of Observation:				5 = defici 10 = profi	
	I t verbal test result qu	iz with pro		selor: PASS	or FAIL (circle one	1	Ciciit
	or passed, continue		-				,	
	,			- J	Score		Comments	
Counseling Sk	ills-Before Rapid Te	est Is Run						
	offered options in t		cedures.					
	carefully explained			ntial				-
results.	, ,	•	.					
3. Counselor	carefully explained	confiden	tial and anony	mous				
testing								
	obtained written in							
	addressed partner							
	gave client subject							
	assessed whether c	lient was	ready to recei	ive				
results tha	it day. ills-While Rapid Tes	at ia Dunni	ing					
	-		<u> </u>					
	identified client's r							
	identified client's sa							
	mainly used non-ju with client.	dgmental	language and	l tone in				
11. Counselor	asked the client ope	en-ended	questions.					
12 Counselor	maintained strong	eve conta	rt and nositiv	e hody				
language.	mamamea strong	cyc conta	ct and positiv	c bouy				
	offered options and	l did not g	ive directives					
Counseling Sk	ills-After Rapid Test	t has Run						
14. Counselor	accurately commun	nicated re	sult to client					
15. Counselor	allowed time for cli	ent to und	lerstand resu	lt.				
16. Counselor prelim. po	made appropriate i	referrals (one to medica	al care if				
	17. Counselor documented and reviewed a risk reduction plan.							
	18. Counselor identified date of last exposure and reviewed the							
	window period, including possible retesting if client was							
	discussed client neo	eds if resu	lt is prelimin	ary				
20. Counselor	20. Counselor accurately completed HIV Test Form-Part 1 (and Part 2 if prelim pos).							
	b Operation Skills							
21. Counselor	set up lab space and	d labeled	devices prope	erly.		T		

e 2015

	_	Revised June				
22. Counselor adhered to all Universal Precautions.						
23. Counselor carefully instructed/demonstrated how to collect specimen and run the test properly.						
24. Counselor did not contaminate specimen or device.						
25. Counselor did move test during processing.						
26. Counselor timed the processing accurately.						
27. Counselor accurately interpreted and documented test result						
28. Counselor recapped all used vials and disposed of used testing supplies in a biohazard container.						
Total Score:						
Scoring Required to Pass: -Each section requires 85% correct to pass, and for those items in bold and under down for each section is as follows: Counseling Skills-Before the Rapid Test is Run = 70 points possible, 60 needed to pass Counseling Skills-While Rapid Test is Running = 60 points possible, 50 needed to pass Rapid Test Lab Operation Skills = 80 points possible, 65 needed to pass Rapid Test Lab Operation Skills = 80 points possible, 65 needed to pass	oass oass	e of 10 (adequate) is required. The break				
Name of person conducting to Name of person conducting to	this observat	tion Counselor #				
Affiliation of Observer to Counselor (i.e. supervisor, regional coordinator)						
Signature and Date of Observer Named Above: Signature		Date				
Write in below the complete physical mailing address where Counselor Certific	cate should b	pe mailed:				
Name of Organization:						
Street Address:City, State,	ZIP:					

Louisiana HIV Prevention Counseling, Rapid Testing and Referral Services Quality Assurance Site Visit Assessment

This form should be completed on the first day of the quality assurance site visit.

SEC'	TION I. Agency Informat	ion Ass	essment Period		
1.	Agency Name				
2.	Name and Title of Supervi	sor/QA Coordinator			
3.	CLIA Waiver NumberExpiration Date				
4.	Is CLIA Waiver displayed properly? Yes No				
5.	Type of Rapid Tests In Use:				
6.	Describe the location when	-			
7. 8.	_	ure Logs Maintained on site? Ye stored testing devices monitored:	s No		
9.	Review the Test Device Temperature Logs for missing entries, days when temperature was out of range, and any corrective actions taken. Record in the table below.				
	Date	Describe Problem/Issue	Describe Action Taken (if any)		
10.	. Describe where Rapid Tes	ting Controls are stored:			
-0.		6 3 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2			

11. Are Rapid Testing Control Logs Maintained on site? Yes No

12. Hov	ow is the temperature of control kits monitored?				
13. Rev	iew the Control K	it Temperature Logs for missing entrie	es, days when temperature was out of range, and	any corrective	
actio	ons taken. Record	I in the table below.			
	Date	Describe Problem/Issue	Describe Action Taken (if any)		
14. Are D	aily Test Logs ma	intained on site? Yes No			
15. How w	vell does the site d	ocument risk reduction plans in client	charts? (review at least 10 charts and indicate wh	at percentage had	
documente	ed risk reduction p	lans)			
16. Are cli	ent files maintaine	ed appropriately? Yes No			
SECTION	II. – Comments/N	Notes/Concerns about rapid testing site			
Use this ren	nainder of this page	and the back if needed to make notes abou	t the site's overall rapid testing policies, any additiona	l concerns, and	
adherences	to SHP protocol.				

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Attachment RT-3.13 (maintain on site)

Risk Reduction Worksheet

Thank you for participating in our HIV & syphilis testing program. If you received a **negative** test result that means the test did not detect any HIV or syphilis antibodies or the p24 antigen in your body. The p24 antigen can take up to 1 month to develop if you've been exposed to HIV, so it's important that you know that you could still be infected with HIV even if you tested negative today, especially if you've been exposed in the last month and your body hasn't developed antibodies yet. So get tested regularly, and at least 1 months after having unprotected sex, injecting drugs, or practicing any other behaviors that could put you at risk for HIV, including coming into contact with any of the 4 bodily fluids that HIV can be transmitted through - blood, semen, vaginal fluid, and breast milk. If you tested **preliminary positive** that means HIV antibodies and/or the p24 antigen were detected by the test, and a confirmatory test is necessary for diagnosis. Please see a medical doctor to learn the best ways to treat the HIV infection; your counselor will help you determine where you might go for medical treatment and can tell you about other types of support available in your area.

During your counseling session today, we talked about behaviors that may put you at risk for HIV and other STDs, and ways to reduce those risks. Below is a summary of your counseling session.

<u>Behaviors</u>	Action Steps to Reduce Risk	Time Frame
Having Anal Sex		
○ w/ condom		
w/out condom		
○ , e.u. ee.u.e		
Having Vaginal Sex		
○ w/ condom		
○ w/out condom		
Having Oral Sex		
○ w/ condom		
○ w/out condom		
Charing Needles or Injection Facilities		
Sharing Needles or Injection Equipment		
Having Unprotected Sex with a Person who is		
HIV+		
○ Other		

Client Signature/Initial	Counselor Number: